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temperature. We have de	termined by simulations how isoled new methods to study how an	epitaxy changes after deposition, if we change lated islands on a solid surface move or shrink ensemble of islands evolves when heated or what understanding of basic phenomena taking plane.	hen
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Final Report, Grant F 49620-98-1-0366 "The Growth of Nanostructures and Composite Films on Solids: Simulations and Phenomenological Theory"

The resources provided by this grant were used for theoretical research in several areas. We performed simulations of a number of phenomena taking place in the early stages of film growth by epitaxy. We have provided a complete description of how the islands formed on the surface during deposition, move[3], evaporate[9], and change shape[4]. We developed a method to simulate coarsening of these islands on a very long time and space scale[5,7]. We proposed and documented by simulations[8,10] that nucleation and growth can be controlled by imposing a periodic strain on the substrate, which will lead to the formation of a periodic array of islands of equal size. To achieve this kind of control has been one of the main goals of nano-science. It has been observed that islands grown on a surface prefer a specific size. We have shown[1] that this may happen because the island size fits a natural length of the electronic wave-function.

In the second part of the grant's duration, we switched to studying various catalytic systems. We have shown that surface strain modifies very substantially the adsorption isotherm[11] and this may explain why ultrasound passing through a catalyst changes its activity. Our current work examines the electronic properties of the γ-alumina[13] (which is the support of choice in most catalysts). This is the first calculation on this very demanding system. We have also determined how Pd atoms migrate on γ-alumina surface[14]. The adsorption of a metal on the surface of another metal, can modify its properties and we looked[16] at how the capacity of adsorbing CO is modified when Fe, Cu, or Ni islands are adsorbed on Ag. We were prompted to do this by very elegant single-molecule experiments by Wilson Ho, whose conclusions are confirmed by our calculations. Finally, we developed a new method for deriving potential expressions from *ab initio* energy calculations[2], proposed a new numerical method for performing quantum dynamics[15], developed an algorithm for solving Maxwell equations and used it to examine how the electromagnetic field under an STM tip is enhanced[12], and wrote a review article of our work on rate theory in quantum systems.

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 $\gamma_{k} = \gamma_{k} \cdot \gamma$

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